

*****STN Columbus *****

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index bioscience

FILE 'DRUGMONOG' ACCESS NOT AUTHORIZED
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INDEX: ADISALERTS, ADISINSIGHT, AGRICOLA,
AIDSLINE, ANABSTR, AQUASCI,
BIOBUSINESS, BIOCOMMERCE, BIOSIS,
BIOTECHABS, BIOTECHDS, BIOTECHNO, CABA,
CANCERLIT, CAPLUS, CEABA, CEN, CIN,
CONFSCI, CROPB, CROPU, DDFB, DDFU,
DGENE, DRUGB, DRUGLAUNCH,
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2000

57 FILES IN THE FILE LIST IN STNINDEX

Enter SET DETAIL ON to see search term postings or to
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=> s (mercaptotripropionamide or phosphonic(l)hair and
(inhib? or depilat? or prevent?)

1 FILE BIOSIS
11 FILES SEARCHED...
1 FILE CABA
9 FILE CAPLUS
20 FILES SEARCHED...
29 FILES SEARCHED...
1 FILE EMBASE
2 FILE IFIPAT
39 FILES SEARCHED...
1 FILE MEDLINE
1 FILE SCISEARCH
52 FILES SEARCHED...
1 FILE TOXLINE
1 FILE TOXLIT
206 FILE USPATFULL
1 FILE WPIDS
56 FILES SEARCHED...

1 FILE WPINDEX

12 FILES HAVE ONE OR MORE ANSWERS, 57
FILES SEARCHED IN STNINDEX

L1 QUE (MERCAPTOTRIPROPIONAMIDE OR
PHOSPHONIC(L) HAIR AND (INHIB? OR DEPILAT? O

R PREVENT?)

=> d rank

F1	206	USPATFULL
F2	9	CAPLUS
F3	2	IFIPAT
F4	1	BIOSIS
F5	1	CABA
F6	1	EMBASE
F7	1	MEDLINE
F8	1	SCISEARCH
F9	1	TOXLINE
F10	1	TOXLIT
F11	1	WPIDS
F12	1	WPINDEX

= file f2-11

COST IN U.S. DOLLARS	SINCE FILE
TOTAL	

ENTRY	SESSION
FULL ESTIMATED COST	2.25
2.40	

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2000

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=> s 11

6 FILES SEARCHED...
L2 19 L1

=> dup rem 12

PROCESSING COMPLETED FOR L2

L3 12 DUPLICATES REMOVED)
ANSWERS '1-9' FROM FILE CAPLUS
ANSWERS '10-11' FROM FILE IFIPAT
ANSWER '12' FROM FILE WPIDS

=> d bib ab 1-12

L3 ANSWER 1 OF 12 CAPLUS COPYRIGHT 2000
ACS DUPLICATE 1
AN 1992:486991 CAPLUS
DN 117:86991

T1 Production of the phytoalexin glyceollin I by soybean
roots in response to
symbiotic and pathogenic infection
AU Schmidt, Petra E.; Parniske, M.; Werner, D
CS Fachbereich Biol., Philipps-Universität Marburg, Germany
SO Bot. Acta (1992), 105(1), 18-25
CODEN: BOACEJ; ISSN: 0932-8629

DT Journal
LA English

AB The amt. of the phytoalexin glyceollin I (I) in root
exudate and root

hairs of individual seedlings of Glycine max was

- analyzed using a R1A. *Bradyrhizobium japonicum* 110spc4, which is able to form N fixing nodules with this plant, caused an increase of lincocq 50-fold in I levels in root exudate relative to uninfected control seedlings. Max. I levels were reached within 10 h of incubation. Elevated I levels were also obsd after incubation of soybean roots in sterile bacterial supernatant, a suspension of autoclaved bacteria or the supernatant from broken cells of *B. japonicum*. Increased I prodn. is not due to the process of active root ***hair*** penetration by the microsymbiont since living bacterial cells are not necessary for the induction. The obsd. I prodn. in *japonicum* is several times lower than that after pathogenic infection. Infection with zoospores of the phytopathogenic oomycete, *Phytophthora megasperma* f. sp. *glyciniae* race 1, leads within 20 h to an accumulation of 7 nmol I seedling in the root exudate of the compatible cultivar Kenwood and 48 nmol -I seedlings in that of the incompatible cultivar Maple Arrow. Apparently, phytoalexins are implicated in detn. of compatibility in pathogenic interactions. Crude cell exts. of different symbiotic bacteria (*B. japonicum* 110spc4, R. meliloti 2011, R. leguminosarum PRE 8, *Sinorhizobium fredii* HH 103) were found to induce different amts. of I in the root exudate. The obsd. I levels could not be correlated with the ability of these rhizobial strains to nodulate soybean. ***inhibition*** of flavonoid and phytoalexin synthesis by (R)-(1-amino-2-phenylethyl) ***phosphonic*** acid (APEP), a specific ***inhibitor*** of the phenylalanine-aminolase (PAL), during the first 20 h of the symbiotic interaction dramatically decreased the no. of nodules formed in root regions that had been in contact with the ***inhibitor***. This effect was obsd. at concns. that
- ***inhibited*** neither bacterial nor plant growth. The implications of these findings for the process of nodule initiation are discussed.
- I3 ANSWER 2 OF 12 CAPLUS COPYRIGHT 2000 ACS DUPLICATE 2
AN 1986:564623 CAPLUS
DN 105:164623
TI Amelioration of cisplatin-induced ototoxicity by fosfomycin
AU Schweitzer, Vanessa G.; Dolan, David F.; Davidson Thomas, Abrams, Gerald
E.; Snyder, Ronald
CS Dep. Otolaryngol. Head Neck Surg., Henry Ford Hosp., Detroit, MI, 48202, USA
SO Laryngoscope (1986), 96(9, pt. 1), 948-58
CODEN: LARYA8; ISSN: 0023-852X
DT Journal
LA English
AB The continued chemotherapeutic application of cisplatin [15663-27-1] necessitates redefn. of its dose-limiting toxicity without decreasing its tumoricidal effect. This research project evaluated the efficacy of fosfomycin [23155-02-4], a ***phosphonic*** acid antibiotic, in decreasing or ameliorating the ototoxicity (high-frequency sensorineural hearing loss) and nephrotoxicity (renal tubular necrosis and interstitial nephritis) of cisplatin. The efficacy of fosfomycin in blocking Pt-induced toxicity in the guinea pig was evaluated histol. and functionally using cyocochleog. and light microscopy of the organ of Corti and the auditory brain stem evoked response (ABR), and light microscopy of renal corticomedullary tissues, small bowel, liver, lung, and peripheral nerve. The results demonstrate that fosfomycin ameliorates the acute renal tubular necrosis and interstitial nephritis and markedly ***inhibits*** the elevation of ABR thresholds and simultaneous outer ***hair*** cell loss that can result from cisplatin administration.
- Fosfomycin should be considered a potential antidote for the dose-limiting ototoxicity and nephrotoxicity of cisplatin chemotherapy.
- I3 ANSWER 3 OF 12 CAPLUS COPYRIGHT 2000 ACS
AN 2000:401835 CAPLUS
DN 133:43508
TI Thienopyran derivatives with activity as potassium channel activators and their use as antihypertensives, antiasthmatics, and hair growth modulators
IN Esch, Peter; Rovinsky, Franz; Towart, Robertson; Christoph, Thomas; Hartmann, Michael; Kealey, George Terence Evelyn
PA Cambridge Bioclinical Limited, UK
SO PCT Int. Appl., 94 pp.
CODEN: PIXXD2
DT Patent
LA English
FAN CNT 1
PATENT NO. KIND DATE APPLICATION
NO. DATE
PI WO 2000/34287 A2 20000615 WO
1999-GB4037 19991203
W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PR: AI GB 1998:26830 19981204
GB 1999:9792 19990428
OS MARPAT 133:43508
AB Novel thienopyran compds. I-III [X H or electron-withdrawing group: Y - spiro fusion with an (un)substituted, unsatd. or monounsatd., alicyclic or heterocyclic ring; Z = H, OH, or alkoxy] compds. having

alkoxy groups), which are useful as reducing agents for hair preps, such as primary agents for wave-setting or straightening curly hair.

pretreatment agents for dyeing hair, oxidative hair dyes, or ***depilatory*** agents, are prepd. This compds. give little damage to hair, reduced odor and flaking, excellent waving effect, and long-lasting wave and are safely used. Thus, 82.5 g monoethanolamine and 25.0 g Et l-cysteinate hydrochloride were placed in a reaction bath, heated at 7-5 mmHg and room temp. for 1 h, and treated with a soln. of 5.4 g NaOH in 20 H₂O under cooling. Excess monoethanolamine was distilled off at 7-5 mmHg and 58-71 degree, to give an oil, which was treated with 200 mL ethanol, refluxed, filtered to remove insol. NaCl, cooled, and filtered to remove pptd. crystals to give 45% (R)-HSCl₂CH(NH₂)CONHCH₂CH₂OH.

L3. ANSWER 7 OF 12 CAPLUS COPYRIGHT 2000
ACS
AN 1996:315324 CAPLUS
DN 124:352330
TI Anti-dandruff hair rinse containing cationic germicide, ammonium conditioner, and metal chelator
IN Hioki, Yuichi; Moriama, Tadashi; Tamura, Yoshinori; Okamoto, Juri; Takeshige, Yuichi
PA Kao Corp., Japan
SO Ger. Offen., 13 pp.
CODEN: GWXXBX
DT Patent
LA German
FAN CNT 1
PATENT NO. KIND DATE APPLICATION
NO. DATE
PI DE 19536420 A1 19960411 DE
1995:19536420 19950929
JP 08099841 A2 19960416 JP 1994:239981
19941004
CN 1126585 A 19960717 CN 1995:117383
19950929
PR:AI JP 1994:239981 19941004
OS MARPAT 124:352330

AB A hair rinse contg. (a) an alkylbenzyltrimethylammonium germicide, (b) a quaternary ammonium-type cationic polymer or cationic surfactant as conditioner, and (c) a chelating agent in a molar ratio to the other 2 components of g:oreq 0.5 shows good conditioning, anti-dandruff, antipruritic, and deodorant activity even in the presence of anionic surfactants. Thus, a hair rinse was prepd. contg. (2-dodecylhexadecyltrimethylammonium chloride 1.5, benzalkonium chloride 1.0, di-Na EDTA 2.0, cetostearyl alc. 3.0, liq. paraffin 1.0, dimethylpolysiloxane 1.0, hydroxyethylcellulose 0.5, methylparaben 0.5, perfume 0.4, and water to 100.0%.

L3. ANSWER 8 OF 12 CAPLUS COPYRIGHT 2000
ACS
AN 1994:143672 CAPLUS
DN 120:143672
TI Composition for treating keratinous fibers containing metal compounds and acids
IN Hirano, Yuji; Kure, Naohisa
PA Kao Corp., Japan
SO PCT Int. Appl., 22 pp.
CODEN: PINXD2
DT Patent
LA English
FAN CNT 1
PATENT NO. KIND DATE APPLICATION
NO. DATE
PI WO 9400099 A1 19940106 WO 1993:JP868
19930625
W: US
RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT,
LU, MC, NL, PT, SE
JP 06009347 A2 19940118 JP 1992:170911
19920629
FP 601163 A1 19940615 EP 1993:013584
19930625
FP 601163 B1 19960320
R: DE, FR, GB
US 5472697 A 19951205 US 1994:182122
19940203
PRAI JP 1992:170911 19920629
WO 1993:JP868 19930625

AB A compn. for treating keratinous fibers comprises: (a) a first agent contg. a metal ion, (b) a second agent contg. an org. or inorg. compd. which is capable of readily permeating into the keratinous fibers and can form water-insol. or sparingly sol. complex together with the metal ion of component (a), and an org. compd. which cannot readily permeate into the keratinous fibers, and which reacts with the metal ion of component (a) to form a water-sol. complex. When the compn. is used for treating keratinous fibers, it allows water-insol. or sparingly sol. complex products to deposit inside the keratinous fibers while ***preventing*** them from depositing on the surface of the keratinous fibers. Because of this mechanism, the present compn. is capable of imparting sufficient firmness and elasticity to keratinous fibers without inducing any objectionable rough or frictional feel to the touch of the hair. For example, a first agent contained hydroxyethyl cellulose 1.3, CaCl₂ 1.5%, and water balance and a second agent contained hydroxyethyl cellulose 1.3, oxalic acid 2.0, 1-hydroxyethylidene-1,1-disulfonic acid 1.0%, arginine q.s., and water balance.

L3. ANSWER 9 OF 12 CAPLUS COPYRIGHT 2000
ACS
AN 1971:532922 CAPLUS
DN 75:132922
TI Aminopolysphonic acids and polyphosphonic acids and their derivatives for the protection of hair
IN Berth, Peter; Reese, Guenter
PA Therachemie Chemisch Therapeutische G.m.b.H.
SO U.S., 3 pp.
CODEN: USXXAM
DT Patent
LA English
FAN CNT 1
PATENT NO. KIND DATE APPLICATION
NO. DATE

PI US 3542918 A 19701124 US 1966-586955
 19661017
 NI 6615210 A 19670525 NI 1966-15210
 19661027
 NI 160161 B 19790515
 BE 689986 A 19670522 BF 1966-89986
 19661121
 IT 1013002 A 19770330 IT 1966-43047
 19661122
 ES 333706 A2 19680101 ES 1966-333706
 19661123
 CH 482444 A 19691215 CH 1966-482444
 19661123
 DK 116529 B 19700119 DK 1966-6068
 19661123
 NO 122856 B 19710823 NO 1966-165707
 19661123
 FI 45213 B 19711231 FI 1966-3108
 19661124
 SE 343475 B 19720313 SE 1966-16110
 19661124
 PR DE 1965-129843 19651124
 AB Damage to human hair during bleaching or dyeing was
 prevented by
 using 0.01-10% of an aminopolysphosphonic acid or deriv.
 alone or in
 combination with a polysphosphonic acid. For example, a
 6 H2O2 bleaching
 soln. (pH 10) was mixed with 0.1%
 aminotris(methylenephosphonic acid) (I)
 or a mixt. of 0.1% of equal parts ethanol-1,
 1-diphosphonic acid and I and
 applied to human hair, and the hair developed a high gloss
 and sheen, with
 little damage. Other P compds. used were
 ethylenediaminetetrakis(methylene
 phosphonic acid), Na, K, or monoethanolamine salt,
 monoethyl
 aminoethane-1, 1-diphosphonate,
 aminotris(methylenephosphonic acid) or
 NH4+ salt, aminotris(isopropylmethylene phosphonic acid) or I
 NH4+ salt.

L3 ANSWER 10 OF 12 IFIPAT COPYRIGHT 2000 IFI
 AN 2699838 IFIPAT:IFIUDB:IFICDB
 TI COMBINED TWO-PART REDUCING
 AGENT HUMECTANT SHAVING SYSTEM FOR
 IMPROVED
 SHAVING COMFORT, BREAKING DISULFIDE
 BONDS, TOILETRIES
 INF Stife, Charles W., New Market, MD

Stoner, Karla L., Frederick, MD
 IN Stife Charles W., Stoner Karla L.
 PAF The Gillette Company, Boston, MA
 PA Gillette Co The (346%)
 ENNAM Page, Thurman K
 AG Williams, Stephan P
 PI US 5500210 19960319 (CITED IN 002 LATER
 PATENTS)
 AI US 1994-247915 19940523
 XPD 23 May 2014
 FI US 5500210 19960319
 DT UTILITY
 FS CHEMICAL
 MRN 007091 MFN: 0351
 CLMN 10
 AB The present invention relates to a method of improving
 shaving comfort by
 softening the hair to be shaved so as to reduce the cutting
 force
 required to cut it. The novel method comprises carrying
 out the following
 sequential steps: (a) contacting an area of hair to be
 shaved with a
 reducing agent that breaks disulfide linkages in hair; (b)
 contacting the
 area of hair treated in step (a) with a humectant and
 allowing it to dry,
 or partially dry; (c) contacting the area treated in step (b)
 with water
 to hydrate the hair; and (d) shaving the hydrated hair of
 step (c).

L3 ANSWER 11 OF 12 IFIPAT COPYRIGHT 2000 IFI
 AN 0252997 IFIPAT:IFIUDB:IFICDB
 TI PHOSPHONIC ACID DERIVATIVES FOR
 PROTECTION OF HAIR FROM DAMAGE IN
 BLEACHING AND DYEING THE SAME
 IN BERTH PETER (DE); REESE GUNTER (DE)
 PA THERACHEMIE CHEMISCH THERAPEUTISCHE
 GESELLSCHAFT MBH GER (84160)
 PI US 3202579 19650824 (CITED IN 018 LATER
 PATENTS)
 AI US 1963-310165 19630919
 XPD 24 Aug 1982
 PRAI DE 1963-T23551 19630302
 FI US 3202579 19650824
 FR 1393604
 DT UTILITY
 FS CHEMICAL
 OS CA 63-4092

L3 ANSWER 12 OF 12 WPIDS COPYRIGHT 2000
 DERWENT INFORMATION LTD
 AN 1994-147833 [18] WPIDS
 DNC C1994-067894
 TI ***Hair*** washing compsn for time-lapse
 colouration
 prevention - contg. anionic surfactant(s), alkanol
 amide(s), amido
 amine type amphoteric surfactant(s), di
 phosphonic acid(s), and
 l-menthol and or dl-menthol.
 DC A96 D21 E19
 PA (SUNZ) SUNSTAR CHEM IND
 CYC 1
 PI JP 06092826 A 19940405 (199418)* 6p
 ADT JP 06092826 A JP 1992-268087 19920909
 PRAI JP 1992-268087 19920909
 AB JP 06092826 A UPAB: 19940622
 The compsn. contains (A) 1-20 wt.% of an anionic
 surfactant(s) of formula
 R1-O-(CH2CH2O)n-SO3M (I), (B) 0.1-10 wt.% of an
 alkanol amide(s) of
 formula (II), (C) 0.1-2.0 wt.% of an amido amine type
 amphoteric
 surfactant(s) of formula (III), (D) 0.001-1 wt.% of
 diphosphonic acid(s)
 of formula (IV), (E) 0.01-3 wt.% of l- and or dl-menthol
 and (F) 0.01-1
 wt.% of an organic acid(s). R1 is 8-20 C alkyl or alkenyl;
 n is 0-5; and M
 is NH4 or cation derived from an alkanolamine. R2 is
 7-21C alkyl or
 alkenyl. R3 and R5 are 7-19C alkyl or alkenyl. R4 and
 R6 are -CH2COOM1 or
 -CH2CH2COOM1; R7 is H, -CH2COOM1 or
 -CH2CH2COOM1; and M1 is H, alkali
 metal or cation derived from an alkanolamine. M2 - M5
 are H or alkali
 metal.
 USE - The compsn. does not cause time-lapse
 colouration.
 Dwg.0 0
 => log y
 COST IN U.S. DOLLARS SINCE FILE
 TOTAL ENTRY SESSION
 FULL ESTIMATED COST 76.09

78.49

DISCOUNT AMOUNTS (FOR QUALIFYING
ACCOUNTS) SINCE FILE TOTAL
ENTRY SESSION
CASH SUBSCRIBER PRICE -5.01
-5.01

STN INTERNATIONAL LOGOFF AT 13:57:11 ON 31
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